II. REMARKS

Formal Matters

Claims 1, 4-8, and 11-35 are pending in this application.

Claims 1, 4-8, and 11 were examined and were rejected. Claims 12-35 were withdrawn from consideration.

Applicants respectfully request reconsideration of the application in view of the remarks made herein.

Rejection under 35 U.S.C.§112, first paragraph

Claims 1, 4-8, and 11 were rejected under 35 U.S.C.§112, first paragraph, as allegedly failing to comply with the enablement requirement.

The Office Action stated that the instant specification has not shown any *in vivo* working examples wherein subjects suffering from hyperlipidemia present reduced severity of symptoms after the administration of an antisense nucleic acid targeting apoE3. Applicants respectfully traverse the rejection.

To comply with 35 U.S.C. § 112, first paragraph, a specification need only enable a skilled artisan to make and use the claimed invention without undue experimentation. Accordingly, a specification complies with the statute even if a reasonable amount of experimentation is required, as long as the experimentation is not "undue."

Applicants respectfully submit that when evaluated in view of the relevant *Wands* factors, the specification clearly enables one of skill in the art to practice the subject invention without undue experimentation. In other words, claims 1, 4-8, and 11 recite subject matter that is adequately described in the specification in such a way as to teach a skilled artisan how to make and use the claimed invention without having to practice undue experimentation.

To aid in determinations of enablement, courts have identified eight factors for consideration: (a) the quantity of experimentation necessary; (b) the amount of direction or guidance presented; (c) the presence or absence of working examples; (d) the nature of the invention; (e) the state of the prior art; (f) the relative skill of those in the art; (g) the predictability or unpredictability of the art; and (h) the breadth of the claims.

¹ Ex Parte Forman., 230 USPQ 546, 547 (Bd.Pat.App & Interf. 1986); and, In re Wands, 8 USPQ2d 1400, 1404 (Fed. Cir. 1988).

(1) the breadth of the claims

The instant claims recite administering an agent that reduces the amount of plasma active apoE in a host. The instant claims encompass the use of antisense nucleic acids that have the ability to effectively reduce expression of apoE, thereby reducing the amount of plasma active apoE, which in turn reduces VLDL production. In order to fall within a claim, an antisense nucleic acid must be able to reduce apoE expression. The number of "agents" encompassed by the claims is thus relatively small.

(2) the state of the art

As of the April 12, 1999 priority date of the instant application, the state of the art of making and using antisense nucleic acids to modulate gene expression was such that a person skilled in the art, given the nucleotide sequence of a given gene, could readily design, make, and use antisense nucleic acids that would reduce the expression of the gene.

The fact that design and use of antisense nucleic acids was a well-developed technology as of the April 12, 1999 priority date is supported by numerous publications in the field. For example, the instant specification cites Wagner et al. ((1996) *Nature Biotechnol*. 14:840-844) which discusses potent and selective inhibition of gene expression by antisense heptanucleotides. Numerous additional papers published before April 12, 1999 attest to the fact that design and use of antisense nucleic acids was a well-developed technology. Numerous textbooks published before April 12, 1999 further demonstrate the fact that the state of the art of antisense technology was highly developed. Such textbooks include, e.g.,:

- 1) Applied Antisense Oligonucleotide Technology (1998) C.A. Stein et al., Eds., Wiley-Liss;
- Clinical Trials of Genetic Therapy with Antisense DNA and DNA Vectors (1998) E. Wickstrom, Ed.,
 CRC;
- 3) Antisense From Technology to Therapy (1997) R. Schlingensiepen, Ed., Blackwell Science;
- 4) Antisense Oligodeoxynucleotides and Antisense RNA: Novel Pharmacological and Therapeutic Agents (1997) B. Weiss, Ed., CRC Press;
- 5) Delivery Strategies of Antisense Oligonucleotide Therapeutics (1995) S. Akhtar, Ed., CRC Press; and
- 6) Antisense Research and Applications (1993) S.T. Crooke and B. Lebleu, Eds., CRC.

Thus, as of the April 12, 1999 priority date, antisense technology was well developed.

(3) the predictability or unpredictability of the art

In making this rejection, the Office Action cited Mercola et al. ((1995) Cancer Gene Therapy 2:47-59; "Mercola"). The Office Action stated that Mercola includes "cautionary remarks concerning the prospects of the antisense gene therapy." Office Action, page 5.

The Office Action quoted Mercola as stating that "as with any therapeutic modality, problems arise, notable: (a) degradation of the oligomer...(b) inefficient cell uptake; (c) nonspecific binding; (d) nonspecific cleavage of mRNA ..." While there may be instances in which some non-functional antisense nucleic acids are generated during the course of experimentation, the courts have clearly taught that even in unpredictable arts the specification does not have to disclose every species of a genus that would work and every species that would not work. Furthermore, the quoted statement from Mercola indicates that the cited problems are nothing more than would be expected to be faced with any therapeutic modality, and as such, are not indicative of problems unique to antisense technology.

Indeed, Mercola states:

"Promising results make the considerable efforts of applying oligodeoxynucleotides in whole animals and in clinical trials more plausible. Conversely, oligodeoxynucleotide experiments which yield promising results in tissue culture can be generalized to the in vivo setting by development of clones of cells bearing plasmid-derived antisense RNA against the same target." Mercola, Abstract.

Mercola indicates that results in tissue culture can be generalized to *in vivo* results, and supports the fact that design and use of antisense nucleic acids was readily accomplished by the person of ordinary skill in the art. Thus, if anything, Mercola supports the fact that the instant claims are enabled.

(4) the quantity of experimentation necessary

The Office Action stated that Branch ((1998) TIBS 23:45-50) "notes the importance of the time and expense necessary to screen large numbers of potential antisense molecules and to carefully monitor their *in vivo* effects, due to non-antisense effects and limits of specificity as well as accessibility of the antisense molecules." Office Action, page 5.

However, the courts have clearly taught that the fact that experimentation may be complex does not necessarily make it undue, if the art typically engages in such experimentation. For example, see MPEP \$2164.01.²

As the court explained³:

"[A] considerable amount of experimentation is permissible, if it is merely routine, or if the specification in question provides a reasonable amount of guidance with respect to the direction in which the experimentation should proceed."

Practitioners in the chemical and molecular biology arts frequently engage in extensive modification of

² See also In re Certain Limited-Charge Cell Culture Microcarriers, 221 USPQ 1165, 1174 (Int'l Trade Comm'n 1983), aff'd sub nom., Massachusetts Institute of Technology v. A.B. Fortia, 227 USPQ 428 (Fed. Cir. 1985).

³ In re Wands 8 USPQ 2d at 1404

reaction conditions and complex and lengthy experimentation where many factors must be varied to succeed in performing an experiment or in producing a desired result. The Federal Circuit has found that such extensive experimentation is not undue in the molecular biology arts. For example, the court concluded that extensive screening experiments, while being voluminous, were not undue in view of the art which routinely performs such long experiments.⁴

The instant claims recite use of an antisense nucleic acid that reduces the amount of plasma active apoE by reducing the expression of apoE. The only experiments, if any, that need be performed to enable the entire scope of the claim are those designed to determine which antisense nucleic acids retain the ability to reduce expression of apoE. Such antisense nucleic acids are determined through routine experimentation, typically employing nothing more than performing techniques that were routine in the art, as noted above. Since these experiments are routine in nature, no undue experimentation is required. In other words, the only experimentation that may be required to enable the claimed invention are those experiments to determine the presence of a certain activity, and since this only requires routine assays, no undue experimentation is necessary.

(5) the relative skill of those in the art

The relevant ordinarily skilled artisan is generally a skilled laboratory technician with experience in molecular biology and/or a scientist with the equivalent of a doctoral degree in molecular biology techniques. Furthermore, such artisans are required to keep abreast of the latest technology through continuing education and reading of scientific journal articles. As such, the skill level of those designing and using antisense nucleic acids assays was high as of the April 12, 1999 priority date.

Furthermore, as noted above, the numerous papers and textbooks in the field of antisense technology attest to the fact that the skill level of those in the field of antisense technology was high as of the April 12, 1999 priority date.

(6) the amount of direction or guidance presented

The instant application provides data showing that overexpression and accumulation of apoE causes hypertriglyceridemia by stimulating VLDL production and by impairing VLDL lipolysis. It follows that reducing the plasma level of active apoE will also reduce the plasma level of VLDL.

Nucleotide sequences of apoE mRNAs were known as of the April 12, 1999 priority date of the instant application and available to the public. See, e.g., GenBank Accession No. M12529 – apoE mRNA sequence, published August 8, 1995; Breslow et al. (1982) *J. Biol. Chem.* 257:14639-14641 – human apoE cDNA sequence; and McLean et al. (1984) *J. Biol. Chem.* 259:6498-6504 – human apoE3 cDNA sequence. Those skilled in the

⁴ Hybritech v. Monoclonal Antibodies, Inc. 231 USPQ 81 (Fed. Cir. 1986)

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art, given the known sequences of apoE genes, and given the advanced state of antisense technology, could have readily made and used antisense nucleic acids that would reduce apoE expression.

(7) the presence or absence of working examples:

Compliance with the enablement requirement under 35 U.S.C. §112, first paragraph, does not require or mandate that a specific example be disclosed. "Nothing more than objective enablement is required, and therefore it is irrelevant whether [a] teaching is provided through broad terminology or illustrative examples."

In sum, the amount of experimentation required to make and use antisense nucleic acid that reduces expression of apoE would not be undue one of skill in the art would be able to perform the experiments as a matter of routine to determine the active nucleic acids, and .

Conclusion as to the rejection under 35 U.S.C. §112, first paragraph

Applicants submit that the rejection of claims 1, 4-8, and 11 under 35 U.S.C.§112, first paragraph, has been adequately addressed in view of the remarks set forth above. The Examiner is thus respectfully requested to withdraw the rejection.

⁵ In re Robins 166 USPQ 552 at 555 (CCPA 1970).

III. CONCLUSION

Applicants submit that all of the claims are in condition for allowance, which action is requested. If the Examiner finds that a telephone conference would expedite the prosecution of this application, the Examiner is invited to telephone the undersigned at the number provided.

By:

The Commissioner is hereby authorized to charge any underpayment of fees associated with this communication, including any necessary fees for extensions of time, or credit any overpayment to Deposit Account No. 50-0815, order number UCAL-121.

Respectfully submitted,

BOZICEVIC, FIELD & FRANCIS LLP

Date: Nov. 22, 2006

Paula A. Borden

Registration No. 42,344

BOZICEVIC, FIELD & FRANCIS LLP 1900 University Avenue, Suite 200 East Palo Alto, CA 94303

Telephone: (650) 327-3400 Facsimile: (650) 327-3231

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